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- Charged organic polymer microbeads in paper making process.
- in a papermaking process, improved drainage and retention are obtained when ionic, organic microbeads of less than about 1,000 mm in diameter if crosslinked or less about than 60 nm in diameter if noncrosslinked are added either alone or in combination with a high molecular weight organic polymer, and/or polysaccharide. Further addition of alum enhances drainage formation and retention properties in papermaking stock with and without the present of other additives used in papermaking orcosses.

BACKGROUND OF THE INVENTION

In the past decade, the concept of using colloidal silica and bentonite to improve drainage, formation and retention has been introduced to papermaking. Fast drainage and greater retention of fines contribute to 10 lower cost in papermaking and improvements are always being sought. U.S. Patent Nos. 4388,150 and 4,385,961 disclose the use of a two-component binder system comprising a cationic starch and an anionic colloidal, silicic acid soil as a retention and when combined with cellulose fibers in a stock from which is made. Finnish Published Specification Nos. 67,735 and 67,735 refer to cationic polymer retention agent compounds including cationic starch and polyacrylamide as useful in combination with an anionic silica to 10 improve sizing. U.S. Patent No. 4,788,653 discloses the use of cationic colloidal silica sol with an anionic copolymer of acrylic acid and acrylamido to render the paper stock resistant to destruction of its retention and dewatering properties by shear forces in the paper-making process. A coaccervate binder, three component system composed of a cationic starch, an anionic high molecular weight polymer and disperse component system composed of a cationic starch, an anionic high molecular weight polymer and dispersed silica having a particle diameter range from 1 to 50 nm is revealed in U.S. Patent Nos. 4,843,801 and 15 4,750,973.

The above Finish publications also disclose the use of bentonite with cationic starch and polyacrylamides. U.S. Patent No. 4,305,781 discloses a bentonite-type clay in combination with high molecular weight, substantially non-ionic polymers such as polyethylene oxides and polyacrylamide as a retention aid. Later, in U.S. Patent No. 4,753,710, bentonite and a substantially linear, cationic polymer such as cationic acrylic polymers, polyethylene limine, polyamine epichlorohydrin, and diallyl dimethyl armonium chloride are claimed to give an improved combination of retention, drainage, drying and formation.

It is noted that the silica sol and bentonite are inorganic microparticle materials.

Latices of organic microparticles have been used in high concentrations of 30-70 lbs/ton to give "highstrength" paper products such as gasket materials, roofing felt, paperboard and floor felt and in paper with 25 30-70% mineral fillers (U.S. Patent No. 4,445,970). It is stated that latices have not been used in fine papermaking because such latices are sticky and difficult to use on a Fourdrinier machine. The latices of the above and following four patent references were made according to U.S. Patent No. 4,056,501. They are all emulsions of polymers made from styrene, butadiene and vinylbenzyl chloride which polymers are reacted with trimethylamine or dimethyl sulfide to produce an "onium" cation which is called a pH independent structured latex of 50 to 1000 nm in diameter. These structured cationic latices are used at high levels of concentration i.e. 30-200 lbs/ton either alone (U.S. Patent No. 4,178,205) or with an anionic, high molecular weight polymer, (U.S. Patent No. 4,187,142) or with an anionic polymer (U.S. Patent No. 4,189,345) or as both cationic and anionic latices (U.S. Patent No. 4,225,383). These latices are preferably from 60-300 nm in size. It has been found, in accordance with the present invention, that noncrosslinked 35 organic microbeads of this size and larger are not effective. Furthermore, the process of the present invention uses organic microbeads at a level of 0.05 to 20 lbs/ton, preferably 0.10 to 7.5 lbs/ton whereas the microbeads of the proceeding five U.S. Patent are used at 30-200 lbs/ton to give strength to paper products such as gaskets with a very high 30-70% mineral content. This prior art does not contemplate the use of charged organic micro-beads as a drainage and retention aid at the very low levels as required by the 40 present invention.

The use of an organic crossilinked microbead, in papermaking is taught in Japanese Patent Tokkal JP355596/83:1988 and Kami Pulp Giljitat Times, gps 1-5, March 1989 as a dual system of a cationic or anionic organic microbead of 1-100 microns and an anionic, cationic or nonionic acrylamide polymer. The waterswelling type, cationic, polymer particle is a crossilinked homopolymer of 2-methacryloyloxyethyl trimethylammonium chloride or a crosslinked copolymer of 2-methacryloyloxy-ethyl trimethylammonium chloride/acrylamide (60/40 weight percent). The acrylamide polymer is an acrylamide homopolymer or acrylamide hydroylstate of 17 mole percent anion-conversion or a copolymer of acrylamide/2-methacryloyloxyethyl trimethylammoniumchloride (75/25 weight percent). The anionic microbead is an acrylamide-acrylic acid copolymer.

EPO 0273605 teaches the addition of microbeads having a diameter ranging from about 48-97 m and produced from terpolymers of vinyl acetate (84.6), ethyl acrylate (85.4) and acrylic addi (4.5) or methacyloritide (85), buyl acrylate (85) and acrylic addi (3). These polymeric beads are disclosed as added to an LBKP pulp sturry in order to evaluate the resultant paper for sizing degree, paper force enhancement and distincerpatability. These polymer beads tall outside the scope of those used in the spreamt invention in that the ionic content thereof is too small to impart any appreciable improvement in retention and drainage in the papermaking process.

The present invention encompasses crosslinked, lonic, organic, polymeric microbeads of less than about 750 nm in diameter or microbeads of less than about 60 nm in diameter if noncrosslinked and water-

insoluble, as a retention and drainage aid, their use in papermaking processes, and compositions thereof with high molecular weight polymers and/or polysaccharides.

EP 0,202,780 describes the preparation of crosslinked, cationic, polyacrylamide beads by conventional inverse emulsion polymerization techniques. Crosslinking is accomplished by the incorporation of difunctional monomer, such as methylenebisacrylamide, into the polymer chain. This crosslinking technology is well known in the art. The patent teaches that the crosslinked beads are useful as flocculants but are more highly efficient after having been subjected to unusual levels of shearing action in order to render them water-soluble.

Typically, the particle size of polymers prepared by conventional, inverse, water-in-oil, emulsion, polymertaidin processes are limited to the range of 1-5 microns, since no particular advantage in reducing the particle size has hitherto been apparent. The particle size which is achievable in inverse emulsions is determined by the concentration and activity of the surfactant(s) employed and these are customarily chosen on the basis of emulsion stability and economic factors.

The present invention is directed to the use, in papermaking, of cationic and anionic, crosslinked, polymenic, microbeads. Microgeis are made by standard techniques and microlatices are purchased commercially. The polymer microbeads are also prepared by the optimal use of a variety of high activity surfactant or surfactant mixtures to achieve submicron size. The type and concentration of surfactant should be chosen to yield a particle size of less than about 750 nm in diameter and more preferably less than about 300 nm in diameter.

SUMMARY OF THE INVENTION

According to the present invention, there is provided a method of making paper from a aqueous suspension of cellulosic papermaking fibers, whereby improved drainage, retention and formation properties are achieved. The method comprises adding to the suspension, from about 0.05 to 20 lbs/bn of an ionic, organic polymer microbead of less than about 750 nanometers in diameter if crosslinked or a polymeric microbead of less than about 80 min diameter if norcosslinked and insoluble. Additionally, from about 0.05 to about 20 lbs/bn, preferably about 0.1 - 5.0 lbs/bn, of a high molecular weight, hydrophilic ionic organic polymer, and/or from about 1.0 to about 500, preferably about 5.0-30.0, lbs/bn of an ionic 30 polysaccharide, such as starch, preferably of a charge opposite that of the microbead, may be used. The synthetic organic polymer and polysaccharide may also be of opposite charge to each other. The addition of the microbead compositions results in significant increase in fiber retention and improvement in drainage and formation, said lbs/bn being based on the dry weight of the paper furnish solids. The organic polymer microbead smay be either cationic or anionic.

Alum or any other active, soluble aluminum species such as polyhydroxyaluminum chloride and/or sulfate and mixtures thereof have been found to enhance drainage rates and retention if they are incorporated into the furnish when used with the microbead compositions 0.1 to 20 lbs/fon, as alumina, based on the dry weight of paper furnish solids, are exemplary.

The microbeads may be made as microemulsions by a process employing an aqueous solution comprising a cationic or anionic monomer and crossinking agent; an oil comprising a saturated hydrocarbon; and an effective amount of a surfactant sufficient to produce particles of less than about 0.75 micron in unswollen number everage particle size diameter. Microbeads are also made as microgels by procedures described by Ying Huang et al., Makromol. Chem. 188, 273-281 (1985) or may be obtained commercially as microglatices. The term "microbead", as used herein, is meant to include all of these configurations, i.e. see beads per se, microgels and microlatices.

Polymerization of the emulsion may be carried out by adding a polymerization initiator, or by subjecting the emulsion to ultraviolet irradiation. An effective amount of a chain transfer agent may be added to the aqueous solution of the emulsion, so as to control the polymerization. It was surprisingly found that the crosslinked, organic, polymeric microbeads have a high efficiency as retention and drainage aids when their particle size is less than about 750 mm in diameter and preferably less than about 50m nm in diameter and that the noncrosslinked, organic, water-insoluble polymer microbeads have a high efficiency when their size is less than about 60 mm. The efficiency of the crosslinked microbeads at larger size than the noncrosslinked microbeads may be attributed to the small strands or tails that protrude from the main crosslinked onlymer.

DETAILED DESCRIPTION OF THE INVENTION INCLUDING PREFERRED EMBODIMENTS

Using the ionic, organic, crosslinked, polymeric microbeads of a diameter less than about 750 nm or

the noncrosslinked, water-insoluble beads of less than about 60 nm in diameter according to this invention, improved drainage, formation and greater fines and filler retention values are obtained in papermaking processes. These additives may be added, alone or in conjunction with other materials, as discussed below, to a conventional paper making stock such as traditional chemical pulps, for instance, biseached and of unbleached sulphate or sulphite pulp, mechanical pulp such as groundwood, thermomechanical or chemithermomechanical pulp or recycled pulp such as deinked waste and any mixtures thereof. The stock, and the final paper, can be substantially unfilled or filled, with amounts of up to about 50%, based on the dry weight of the stock, or up to about 40%, based on dry weight of paper of filler, being exemplary. When filler is used any conventional filler such as calcium carbonate, clay, titanium dioxide or talc or a combination may be present. The filler, if present, may be incorporated into the stock before or after addition of the microbeads. Other standard paper-making additives such as rosin sizing, synthetic sizings such as alkyl succinic anhydride and alkyl ketene dimer, alum, strength additives, promoters, polymeric coagulants uch as low molecular weight polymers, dye fixatives, etc. and other materials that are desirable in the papermaking process, may also be added.

The order of addition, specific addition points, and furnish modification itself are not critical and normally will be based on practicality and performace for each specific application, as is common papermaking practise.

When using cationic, high molecular weight polymer(s), or polysaccharides, and anionic microbeads, the preferred sequence of addition is cationic, high molecular weight polymer and then anionic bead. 20 However, in some cases the reverse may be used. When a cationic polysaccharide such as starch and a cationic polymer are both used, they can be added separately or together, and in any order. Furthermore, their individual addition may be at more than one point. The anionic microbeads may be added before any cationic components or after them with the latter being the preferred method. Split addition may also be practised. Preferred practise is to add cationic polysaccharide before high molecular weight cationic polymer. The furnish may already have cationic starch, alum, cationic (or anionic or both cationic and anionic) polymers of molecular weight equal or less than 100,000, sodium aluminate, and basic aluminum salts (e.g., polyaluminum chloride and/or sulfate) and their levels may be varied to improve the response of the furnish, as discussed above. Addition points are those typically used with dual retention & drainage systems (pre-fan pump or pre-screen for one component and pre- or post-screens for another). However, adding the last component before the fan pump may be warranted in some cases. Other addition points that are practical can be used if better performance or convenience is obtained. Thick stock addition of one component is also possible, although thin stock addition is preferred. However, thick stock and/or split thick and thin stock addition of cationic starch is routinely practised and these addition modes are applicable with the use of the microbead as well. Addition points will be determined by practicality and by the possible 35 need to put more or less shear on the treated system to ensure good formation.

When using high molecular weight, anionic polymer(s) and cationic microbeads, the preferred sequence is anionic polymer and then cationic beads, although in some cases the reverse may be used. When anionic polymer and anionic polysaccharide are both used, they can be added separately or together, and in any order.

The microbeads may also be used in combination with high molecular weight ionic polymers of similar or opposite charge,

The microbeads are crosslinked, cationic or anionic, polymeric, organic microparticles having an unswollen number average particle size diameter of less than about 750 nanometers and a crosslinking agent content of above about 4 molar parts per million based on the monomeric units present in the 45 polymer and are generally formed by the polymerization of at least one ethylenically unsaturated cationic or anionic monomer and, optionally, at least one non-ionic comonomer in the presence of sald crosslinking agent. They preferably have a solution viscosity (SV) of about 1.1-2.0 mPa.s.

Cationic microbeads used herein include those made by polymertzing such monomers as ciallyldialkylammmonium halides; acryloxyalkyltrimethylammonium chioride; (meth)acrylates of dialkylaminoalkyl compounds, and salts and quaternaries thereof and, monomers of N.N-dialkylaminoalkyl(meth)acrylamides; and salt and quaternaries thereof, such as N.N-dimethyl aminoethylacrylamides; (meth)acrylamideorpoytrimethylammonium chioride and the acid or quaternary salts of N.N-dimethylaminoethylacrylate and the like. Cationic monomers which may be used herein are of the following general formulae:

(I)
$$CH_2 = C - C - X - R - N^* - R_3$$
 Z^-

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where R₁ is hydrogen or methyl, R₂ is hydrogen or lower alkyl of C₁ to C₄, R₃ and/or R₄ are hydrogen, alkyl of C₁ to C₁₂, aryl, or hydroxyethyl and R₂ and R₃ or R₂ and R₄ can combined to form a cyclic ring or containing one or more hetero atoms, Z is the conjugate base of an acid, X is oxygen or -NR₁ wherein R₁ is as defined above, and A is an alkylene group of C₁ to C₁₂; or

where R_5 and R_6 are hydrogen or methyl, R_7 is hydrogen or alkyl of C_1 to C_{12} and R_8 is hydrogen, alkyl of C_1 to C_{12} , benzyl or hydroxyethyl; and Z is as defined above.

Anionic microbeads that are useful herein those made by hydrolyzing acrylamide polymer microbeads etc. those made by polymerizing such monomers as (methyljacrylic acid and their satts, 2-acrylamido-2-methylpropane sulfonate, sulfoethyl-(meth)acrylate, vinylsulfonic acid, styrene sulfonic acid, maleic or other dibasic acids or their satts or mixtures thereof.

Nonionic monomers, suitable for making microbeads as copolymers with the above anionic and cationic monomers, or mixtures thereof, include (meth)acrylamide; N-alkyacrylamides, such as N-methylacrylamide; NN-dialkylacrylamides, such as NN-dimethylacrylamide; methyl acrylate; methyl methacrylate; acrylonitile; N-vinyl methylacetamide; N-vinyl methyl formamide; vinyl acetate; N-vinyl pyrrofidone, mixtures of any of the forecoping and the like.

These ethylenically unsaturated, non-ionic monomers may be copolymerized, as mentioned above, to produce cationic, anionic or amphoteric copolymers. Preferably, acrylamide is copolymerized with an ionic and/or cationic monomer. Cationic or anionic copolymers useful in making microbeads comprise from about 40 0 to about 99 parts, by weight, of non-ionic monomer and from about 100 to about 1 part, by weight, of cationic or anionic monomer, based on the total weight of the anionic or cationic and non-ionic monomers, preferably from about 10 to about 90 parts, by weight, of non-ionic monomer and about 10 to about 90 parts, by weight, of cationic or anionic monomer, same basis i.e. the total ionic charge in the microbead must be greater than about 1%. Mixtures of polymeric microbeads may also be used if the total ionic 45 charge of the mixture is also over about 1%. If the anionic microbead is used alone, i.e. in the absence of high molecular weight polymer or polysaccharide, in the process of the present invention, the total anionic charge thereof must be at least about 5%. Most preferably, the microbeads contain from about 20 to 80 parts, by weight, of non-ionic monomer and about 80 to about 20 parts by weight, same basis, of cationic or anionic monomer or mixture thereof. Polymerization of the monomers occurs in the presence of a 50 polyfunctional crosslinking agent to form the cross-linked microbead. Useful polyfunctional crosslinking agents comprise compounds having either at least two double bounds, a double bond and a reactive group, or two reactive groups. Illustrative of those containing at least two double bounds are N,Nmethylenebisacrylamide: N.N-methylenebismethacrylamide: polyethyleneglycol polyethyleneglycol dimethacrylate; N-vinyl acrylamide; divinylbenzene; triallylommonium salts, N-methylal-55 Ivlacrylamide and the like. Polyfunctional branching agents containing at least one double bond and at least one reactive group include glycidyl acrylate; glycidyl methacrylate; acrolein; methylolacrylamide and the like. Polyfunctional branching agents containing at least two reactive groups include dialdehydes, such as gyloxal; diepoxy compounds; epichlorohydrin and the like.

Crosslinking agents are to be used in sufficient quantities to assure a cross-linked composition. Preferably, at least about 4 molar parts per million of crosslinking agent based on the monomeric units present in the polymer are employed to induce sufficient crosslinking and especially preferred is a crosslinking agent content of from about 4 to about 6000 molar parts per million, most preferably, about 20-5 4000.

The polymeric microbeads of this invention are preferably prepared by polymerization of the monomers in an emulsion as disclosed in application, Serial No. (Attorney Docket 31320), Polymerization in microemulsions and inverse emulsions may be used as is known to those skilled in this art. P. Spelser reported in 1978 and 1977 a process for making spherical "nanoparticles" with diameters less than 800 Å by (1) solubilizing monomers, such as acrylamide and methylenebisacrylamide, in micelles and (2) polymerizing the monomers, See J. Pharm. Sa., 65(12), 1763 (1976) and United States Patient No. 4,021,864. Both inverse water-in-oil and oil-in-water "nanoparticles" were prepared by this process. While not specifically called microemulsion polymerization by the author, this process does contain all the features which are currently used to define microemulsion polymerization. These reports also constitute the first examples of polymerization of acrylamide in a microemulsion. Since then, numerous publications reporting polymerization of hydrophobic monomers in the oil phase of microemulsions have appeared. See, for examples, U.S. Patent Nos. 4,521,317 and 4,681,912; Stoffer and Bone, J. Dispersion Sci. and Tech., 1(1), 37, 1980; and Atik and Thomas, J. Am. Chem. Soc., 103 (14), 4/27 (1981); and 68 2161492A.

The cationic and/or anionic emulsion polymerization process is conducted by (i) preparing a monomer so emulsion by adding an aqueous solution of the monomers to a hydrocarbon liquid containing appropriate surfactant or surfactant initiure to form an inverse monomer emulsion consisting of small aqueous droplets which, when polymerized, result in polymer particles of less than 0.75 micron in size, dispersed in the continuous oil phase and (il) subjecting the monomer intercomulsion to free radical polymerization.

The aqueous phase comprises an aqueous mixture of the cationic and/or anionic monomers and optionally, a non-ionic monomer and the crosslinking agent, as discussed above. The aqueous monomer mixture may also comprise such conventional additives as are desired. For example, the mixture may contain chelating agents to remove polymerization inhibitors, pH adjusters, initiators and other conventional additives.

Essential to the formation of the emulsion, which may be defined as a swollen, transparent and 30 thermodynamically stable emulsion comprising two liquids insoluble in each other and a surfactant, in which the micelles are less than 0.75 micron in diameter, is the selection of appropriate organic phase and surfactant.

The selection of the organic phase has a substantial effect on the minimum surfactant concentration necessary to obtain the inverse emulsion. The organic phase may comprise a hydrocarbon or hydrocarbon mixture. Saturated hydrocarbons or mixtures thereof are the most suitable in order to obtain inexpensive formulations. Typically, the organic phase will comprise benzene, toluene, fuel oil, kerosene, odorless mineral spirits or mixtures of any of the foreocing.

The ratio, by weight, of the amounts of aqueous and hydrocarbon phases is chosen as high as possible, so as to obtain, after polymerization, are mulsion of high polymer content. Practically, this ratio may range, or example for about 0.5 to about 3.1, and usually approximates about 1.1; respectively.

The one or more surfactants are selected in order to obtain HLB (Hydrophilic Lipophilic Balance) value ranging from about 8 to about 11. Outside this range, inverse emulsions are not usually obtained. In addition to the appropriate HLB value, the concentration of surfactant must also be optimized, i.e. sufficient of the many of the option and inverse emulsion. Too low a concentration of surfactant leads to inverse emulsions of the prior art and too high a concentrations results in undue costs. Typical surfactants useful, in addition to those specifically discussed above, may be anionic, cationic or nonlonic and may be selected from polyoxyethylene (20) sorbitan trioleate, sorbitan trioleates, sorbitan trioleates,

Polymerization of the emulsion may be carried out in any manner known to those skilled in the art.
Initiation may be effected with a variety of thermal and redox tree-radical initiators including azo compounds, such as azobisisobutyronitrile; peroxides, such as 1-butyl peroxide; organic compounds, such as potassium persultate and redox couples, such as ferrous ammonium suftate/ammonium persultate. Polymerization may also be effected by photochemical irradiation processes; irradiation, or by ionizing radiation and the self-cate by inversion by adding it to water which may contain a breaker surfactant. Optionally, the polymer may be recovered for the emulsion by stripping or by adding the emulsion to a solvent which precipitates the polymer, e.g. isopropanol, filtering of the resultant solids, dyring and redispersing in water.

The high molecular weight, ionic, synthetic polymers used in the present invention preferably have a

molecular weight in excess of 100,000 and preferably between about 250,000 and 25,000,000. Their anionicity and/or cationicity may range from 1 mole percent to 100 mole percent. The ionic polymer may also comprise homopolymers or copolymers of any of the ionic monomers discussed above with regard to the ionic beads, with acrylamide copolymers being preferred.

The degree of substitution of cationic starches (or other polysaccharides) and other non-synthetic based polymers may be from about 0.01 to about 0.02. Amphoteric starches, preferably but not exclusively with a net cationic starch, may also be used. The degree of substitution of anionic starches (or other polysaccharides) and other non-synthetic-based polymers may be from 0.01 to about 0.7 or greater. The ionic starch may be made from starches derived from any other or other polysaccharides) and other non-synthetic-based polymers may be common starch producing materials, e.g., potato starch, corn starch, wazy maize, etc. For example, a cationic potato starch made by treating potato starch with 3-chloro-2-hydroxypropylitmethylammonium chloride. Mixtures of synthetic polymers and e.g. starches, may be used. Other polysaccharides useful herein include guar, cellulose derivatives such as carboxymethyloselulose and the like.

It is also preferred that the high molecular weight, ionic polymer be of a charge opposite that of the microbead and that if a mixture of synthetic, ionic polymers or starch be used, at least one be of a charge opposite that of the microbead. The microbeads may be used as such or may be replaced in part, i.e. up to about 50%, by weight, with bentonite or a silica such as colloidal silica, modified colloidal silica etc. and still fall within the scope of the percent invention.

The instant invention also relates to compositions of matter comprising mixtures of the above-described oric microbeads, high molecular weight, ionic polymers and polysaccharides. More particularly, compositions comprising a mixture of A) an ionic, organic, polymer microbead or less than about 750 nanometers in diameter if cross-linked and less than 60 nanometers in diameter if non-cross-linked and water-insoluble and B) a high molecular weight ionic polymer, the ratio of A): B) ranging from about 1:4:00 to A) respectively. Additionally, the compositions may contain the microbead A) and C) an ionic polymer (b) ranging from about 2:0:1 to about 1:1:000, respectively. Still further, the compositions are contain the microbead A), the polymer B) and the polysaccharide C), the ratio of A) to B) plus C) ranging from about 20:1:1:000, respectively Adott 1:000 but 1:1:000, respectively.

Paper made by the process described above also constitutes part of the present invention.

The following examples are set forth for purposes of illustration only and are not be construed as ilmitations on the present invention except as set forth in the appended claims. All parts and percentages are by weight unless otherwise specifices.

In the examples which follow, the ionic organic polymer microbead and/or the high molecular weight, ionic polymer and/or ionic starch are added sequentially directly to the stock or just before the stock reaches the headbox.

Unless otherwise specified, a 70/30 hardwood/softwood bleached kraft pulp containing 25% CaCO₃ is used as furnish at a pri of 8.0. Retention is measured in a Britt Dynamic Drainage Jar. First Pass Retention (FPR) is calculated as follows:

FPR = <u>Headbox Consistency</u> - <u>Tray Water Consistency</u> Head Box Consistency

First Pass Retention is a measure of the percent of solids that are retained in the paper. Drainage is a measure of the time required for a certain volume of water to drain through the paper and is here measured as a 10x drainage. (K. Britt, TAPPI 63(4) p67 (1980). Hand sheets are prepared on a Noble and Wood sheet machine.

In all the examples, the ionic polymer and the microbead are added separately to the thin stock and subjected to shear. Except when noted, the charged microbead (or silica or bentonite) is added last. Unless noted, the first of the additives is added to the test turnish in a "Vaned Britt Jat" and subjected to 800 rpm stirring for 30 seconds. Any other additive is then added and also subjected to 800 rpm stirring for 30 seconds. The respective measurements are when carried out.

Doses are given on pounds/ton for fumish solids such as pulp, fillers etc. Polymers are given on a real basis, silica as SiO₂ and starch, clay and bentonite are given on an as is basis.

55 I. Cationic polymers used in the examples are:

Cationic Starch: Potato starch treated with 3-chloro-2-hydroxypropyltrimethylammonlum chloride to give a 0.04 degree of substitution.

10 AETMAC/90 AMD: A linear cationic copolymer of 10 mole % of acryloxyethyltrimethylammonium chloride and 90 mole % of acrylamide of 5.000.000 to 10,000.000 mol. wt. with a charge density of 1.2

5 AETMAC/95 AMD: A linear copolymer of 5 mole % of acryloxyethltrimethylammonium chloride and 90 5 mole % of acrylamide of 5,000,000 to 10,000,000 mol. wt.

55 AETMAC/45 AMD: A linear copolymer of 55 mole % of acryloxyethyltrimethylammonium chloride and 45 mole % of acrylamide of 5,000,000 to 10,000,000 mol. wt. and a charge density of 3.97 meg/g.

40 AETMAC/60 AMD: A linear copolymer of 40 mole % of acryloxyethyltrimethylammonium chloride and 60 mole % of acrylamide of 5,000,000 to 10,000,000 mol. mt.

50 EPI/47 DMA 3 EDA: A copolymer of 50 mole % of epichlorohydrin, 47 mole % of dimethylamine and 3.0 mole % of ethylene diamine of 250,000 mol. wt.

II. Anionic Polymers used in the examples are: 30 AA/70 AMD: A linear copolymer of 30 mole % ammonium acrylate and 70 mole % of acrylamide of 15 15,000,000 to 20,000,000 mol. wt.

7AA/93 AMD: A linear copolymer of 7 mole % ammonium acrylate and 93 mole % of acrylamide of 15.000.000 to 20,000,000 mol. wt.

10 APS/90 AMD: A linear copolymer of 10 mole % of sodium 2-acrylamido-2-methylpropanesulfonate and 90 mole % of acrylamide of 15,000,000 to 20,000,000 mol. wt.

III. Anionic particles used in the examples are:

SILICA: Colloidal silica with an average size of 5 nm, stabilized with alkali and commercially available. BENTONITE: Commercially available anionic swelling bentonite from clays such as sepiolite, attapulgite or 25 montmorillonite as described in U.S. Pat. No. 4,305,781.

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IV. Latices used in the examples are:

30	<u>Latex</u> Polystyrene	Particle <u>Size in rm</u> 98	Charge Density A / Charge Group 1.4 x 10
35	Polystyrene	30	1.1 x 10 ³
	Polystyrene	22	0.36 x 10 ³

V. Microbeads used in the examples are:

30 AA/70 AMD/50 ppm MBA: An inverse emulsion copolymer of 30 mole % of sodium acrylate and 70 mole % of acrylamide crosslinked with 50 ppm of methylenebisacrylamide with a particle diameter of 1,000-45 2,000 nm; SV-1.64 mPa.s.

40 AA/60 MBA: A microbead dispersion of a copolymer of 40 mole % of ammonium acrylate and 60 mole % of N,N'-methylenebisacrylamide (MBA) with a particle diameter of 220 nm.

30 AA/70 AMD/349 ppm MBA: A microemulsion copolymer of 30 mole % of sodium acrylate and 70 mole % of acrylamide crosslinked with 349 ppm of N,N'-methylenebisacrylanide (MBA) of 130 nm particle 50 diameter, SV-1.17 to 1.19 mPa.s

30 AA/70 AMD/749 ppm MBA: A microemulsion copolymer of 30 mole % of sodium acrylate and 70 mole % of acrylamide crosslinked with 749 ppm of N,N'-methylenebisacrylamide (MBA), Sv-1.06 mPa.s.

60 AA/40 AMD/1,381 ppm MBA: A microemulsion copolymer of 60 mole % of sodium acrylate and 40 mole % of acrylamide crosslinked with 1,381 ppm of N,N'-methylene-bis acrylamide (MBA) of 120 nm particle 55 diameter: SV-1.10 mPa.s.

30 APS/70 AMD/995 ppm MBA: A microemulsion copolymer of 30 mole % of sodium 2-acrylamido-2methylpropane sulfonate and 70 mole % of acrylamide cross-linked with 995 ppm of methylenebisacrylamide (MBA); SV-1.37 mPa.s.

30 AA/70 AMD/1000 ppm MBA 2% SURFACTANT (TOTAL EMULSION): A microemulsion copolymer of 30 mole % of sodium acrylate and 70 mole % of acrylamide crosslinked with 1,000 ppm of N,N*-methylenebisacrylamide with 2% diethanolamide oleate and 494*mn particle diameter.

30 AA/70 AMD/1,000 ppm MBA/ 4% SURFACTANT (TOTAL EMULSION): A microemulsion copolymer of 5 mole % of sodium acrylste and 70 mole % of acrylamide crosslinked with 1,000 ppm of N,N-methylenebisacrylamide with 4% diethanolamide oleate and of 149mp particle diameter, SY-102 mPa.s 30 AA/70 AMD/1,000 ppm MBA/8% SURFACTANT(TOTAL EMULSION): A Microemulsion copolymer of 30 mole % of sodium acrylate and 70 mole % of acrylamide crosslinked with 1000 ppm of N,N-methylenebisacrylamide with 8% diethanolamide oleate and of 108mp particle diameter, SY-1.06 mPa.s.

10 * The unswollen number average particle diameter in nanometers is determined by the quasi-elastic light scattering spectroscopy (QELS).

Procedure for the Preparation of Anionic Microemulsions
15 30 AA/70 AMD/349 ppm MBA - 130 nm

An aqueous phase is prepared by sequentially mixing 147 parts of acrylic acid, 200 parts deionized water, 144 parts of 56.7% sodium hydroxide, 343.2 parts of acrylamide crystal, 0.3 part of 10% pents of the parts of the parts of the parts of deionised water, and 1.5 parts of deionised water, and 1.5 parts of deionized water, and 1.5 parts of deionized water, 0.25 part of 1% though parts of the resultant aqueous phase solution, 6.5 parts of deionized water, 0.25 part of 1% though physical parts of the 3.50 parts of 0.61% methylene bisscrylamide are added. 120 Parts of the aqueous phase are then mixed with an oil phase containing 77.8 parts of low odor parefilm oil, 3.6 parts of softbian seequiclostes and 21.4 parts of polycyyethylene sorbitol hexaclested.

This resultant clear, microemulsion is deaerated with nitrogen for 20 minutes. Polymerization is initiated 25 with gaseous SQ₂, allowed to exotherm to 40°C and controlled at 40°C (+ 5°C) with ice water. The ice water is removed when cooling is no longer required. The nitrogen is continued for one hour. The total polymerization time is 2.5 hours.

For purposes of use in the instant process, the polymer may be recovered from the emulsion by stripping or by adding the emulsion to a solvent which precipitates the polymer, e.g. isopropanol, filtering of the resultant solids, and redispersing in water for use in the papermaking process. The precipitated polymer microbeads may be dried before redispersion in water.

Alternatively, the microemulsion per se may also be directly dispersed in water. Depending on the surfactant and levels used in the microemulsion, dispersion in water may require using a high hydrophilic lipopilic balance (HLB) inverting surfactant such as ethoxylated alcohols; polyoxyethlated sorbitol hexacleate; diethanolamine oleate; ethoxylated laurel sulfate et. as in known in the art.

The concentration of the microbeads in the above-described redispersion procedures is similar to that used with other thin stock additives, the initial dispersion being at least 0.1%, by weight. The dispersion may be redilluted 5-10 tidd just before addition to the papermaking process.

40 Preparation of Cationic Organic Microbead 40 AETMAC/60 AMD/100 ppm MBA - 100 nm By microemulsion

An aqueous phase containing 21.3 parts, by weight of acrylamide, 51.7 parts of a 75% acryloxyethyltrinethyl ammonium chloride solution, 0.07 part of 10% diethylenetriamline pentaacotate (penta sodium 45 ast), 0.7 part of 1% t-butyl hydroperoxide and 0.06 part of methylenebiascrylamide dissolved in 65.7 parts of deionized water is prepared. The pH is adjusted to 3.5 (±0.1). An oil phase composed of 8.4 parts of sorbitan sesquideate, 5.16 parts of ployxyethylene sorbitol hexapleate dissolved in 170 parts of a low paraffin oil is prepared. The aqueous and oil phase are mixed together in an air tight polymerization reactor fitted with a nitrogen spareg tube, thermometer and activator addition tube. The resultant clear microemulsion is sparged with nitrogen for 30 minutes and the temperature is adjusted to 27.5°C. Gaseous studied dioxide activator is then added by bubbling nitrogen through a solution of sodium metabisuffite. The polymerization is allowed the oxotherm to its maximum temperature (about 52°C) and then cooled to 25°C.

The particle diameter of the resultant polymer microbead is found to be 100 nm. The unswollen number average particle diameter in nanometers (nm) is determined by quasi-elastic light scattering spectroscopy (QELS). The SV is 1.72 mPa.s.

Preparation of Cationic Organic Inverse Emulsion 40 AETMAC/60 AMD/100 ppm MBA 1,000 nm by Inverse Emulsion Polymerization

An aqueous phase is made by dissolving 87.0 parts of commercial, crystal acrylamide (AMD), 210.7 parts of a 75% acryloxyethytrimethylammonium chloride (AETMAC) solution, 4.1 parts of ammonium sulfate, 4.9 parts of a 5% ethylene diaminetetraacetic acid (disodium sail) solution, 0.245 part (1000 wppm) of methylenebisacrylamide (MBA) and 2.56 parts of t-burlyl hydroperoxide into 189 parts of delonized water. The pH is adjusted to 3.5 (20.1) with sulfuric acid.

The oil phase is made by dissolving 12.0 gms of sorbitan monocleate into 173 parts of a low odor paraffin oil.

The aqueous phase and oil phase are mixed together and homogenized until the particle size is in the 1.0 micron rance.

The emulsion is then transferred to a one liter, three-necked, creased flask equipped with an agitator, nitrogen sparge tube, sodium metabisulfite activator feed line and a thermometer.

The emulsion is agitated, sparged with nitrogen and the temperature adjusted to 25°C. After the emulsion is sparged 30 minutes, 0.3% sodium metabisuffite (MBS) activator solution is added at a 0.028 mi/minute rate. The polymerization is allowed to exotherm and the temperature is controlled with ice water.

When cooling is no longer needed, the 0.8% MBS activator solution/addition rate is increased and a heating mantle is used to maintain the temperature. The total polymerization time takes approximately 4 to 5 hours using 11 mis of MBS activator. The finished emulsion product is then cooled to 25°C.

The particle diameter is found to be 1,000 nm. The unswollen number average particle diameter in nanometers is determined by the quasi-elastic light scattering spectroscopy (QELS). The SV is 1.24 mPa.s.

EXAMPLE 1

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Using the paper-making procedure described above, the drainage times are measured on 1) sikaline stochatining 5% CaCOs, alone, 2) the same stock with added linear, high molecular weight cationic copolymer of 10 mole % acrylcoxyethytimethylamnoration chloride and 90 mole % of acrylamide (10 AETMAC99 AMD) and 3) the same stock with added cationic copolymer and anionic microbead made from 30 mole % acrylic acid 70 mole % of acrylamide (30 AATO AMD) and cross-linked with 349 ppm of methylenebisacrylamide (MBA) of 130 nm particle diameter and added as a redispersed 0.02% aqueous solution. The results are shown in Table I, below.

TABLE I

Cationic Polymerlbs/Ton	Anionic Microbead	Drainage inSeconds	
-0-	-0-	88.4	
-2-	-0-	62.3	
-2-	0.5	37.5	

The addition of cationic polymer reduces drainage time from 88.4 to 62.3 seconds. Surprisingly mines by another 24.8 seconds to 37.5 seconds, a 39.8% reduction which is a significant improvement in drainage times.

45 EXAMPLE 2

The alkaline furnish used in this example contains 5.0 lbs/ton of cationic starch. To this furnish is added to following additives as described in Example 1. Drainage times are then measured and reported in Table II, below.

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TARLE TIT

Cationic Polymer lbs/Ton -0-	Anionic Microbead 1bs/Ton -0-	Drainage in Seconds 121.9
1 - 10 AEIMAC/90 AMD 1 - 10 AEIMAC/90 AMD	-0- 0.5 - 30 AA/70	89.6
	AMD/ 349 ppm = 130 m	57.8

In the presence of a mixture of high molecular weight cationic polymer and, cationic starch, anionic polymer microbeads greatly improves drainage.

EXAMPLE 3

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Following the procedure of Example 1, various other comparative runs are made using a second alkaline stock containing 10 lbs/lon of cationic starch, and bentonite, as disclosed in U.S. Patent 4,753,710, in order to show the benefits of the use of organic microbeads in accordance with the invention hereof. The results are shown in Table III, below.

TABLE III

9	Cationic Polymer	Anionic Micro- Particle (1bs./Ton) -0-	Drainage in Seconds 132.3
	1.0 - 10 AETMAC/90 AMD	5.0 - Bentonite	53.1
5	1.0 - 10 AETMAC/90 AMD	0.5 - 30 AA/70 AMD/ 349 ppm MBA - 130 mm	55.1
	1.0 - 10 AETMAC/90 AMD	0.5 - 100AA-1985 ppm MBA-80 nm	65.1
	1.0 - 55 AETMAC/45 AMD	5.0 - Bentonite	
)	· · · · · · · · · · · · · · · · · · ·		76.4
	1.0 - 55 AETMAC/45 AMD	0.5 - 30 AA/70 AMD/ 349 ppm MBA - 130 mm	55.4
5	1.0 - 55 AETMAC/45 AMD	0.5 - 60 AA/40 AMD/ 1,381 ppm MBA - 120 rm	45.7
	1.0 - 55 AETMAC/45 AMD	0.5 - 100AA-1985 ppm MBA	48.6

When the 10% cationic polymer AETMAC/AMD (10/90) is used in conjunction with 5.0 lbs. of bentonite, similar drainage results to those obtained using only 0.5 lb. of 30% anionic microbead AA/AMD (30/70) in place of the bentonite, are obtained. With a 55% cationicity polymer, bentonite gives a solewar drainage rate of 76.4 seconds and the 30% anionic microbead about the same drainage rate of 55.4 seconds. With the 5 higher cationicity polymer (55%) and 0.5 lbs/ton of a high anionicity microbead, AA/AMD (60/40) a far superior drainage time of 45.7 seconds is obtained, using far less sadditive.

EXAMPLE 4

An alkaline paper stock containing 10 pounds/fon of cattonic starch is treated as described in Example 1. The results are shown in Table IV, below.

		TABLE IV	
5	Cationic Polymer	Anionic Micro- particle 1bs/Ton -0-	Drainage in Seconds 115.8
	0.5 - 10 AETMAC/90 AMD	-0-	83.5
10	0.5 - 10 AEIMAC/90 AMD	5.0 - Bentonite	51.1
	0.5 - 10 AETMAC/90 AMD	0.5 - 30 AA/70 AMD/ 349 ppm MBA - 130 nm	57.3
15	0.5 - 55 AETMAC/45 AMD	0.5 - 60 AA/40 AMD/ 1,381 ppm - 120 mm	46.1
	1.0 - 10 AETMAC/90 AMD	5.0 - Bentonite	42
20	1.0 - 55 AEIMAC/45 AMD	0.5 - 60 AA/40 AMD/ 1,381 ppm EMA - 120 rm	38.9

The combination of 0.5 lb/ton of cationic polymer and 5.0 lbs/ton of bentonite gives a good drainage of 51.5 seconds, somewhat better than the 0.5 lb of 30% anionicity microbeads, i.e. 57.3 seconds. However, bentonite is inferior to the results achieved using 0.5 lb/ton of a higher (60%) anionicity polymer, i.e. of 46.1 seconds. Increasing the amount of cationic polymer to 1.0 lb/ton results in improved bentonite and 60% anionic polymer microbead times of 42 and 38.9 seconds, however, the microbead results are again superior.

30 EXAMPLE 5

The procedure of Example 1 is again followed except that first pass retention values are measured. The organic anionic microbead is compared at a 0.5 lbs/fon rate to 2.0 lbs/fon of sitica and 5.0 lbs/fon of bentonite in an alkaline paper stock as known in the art. The organic, 30% anionic polymer microbeads give the best retention values at a lower concentration, as shown in Table V, below.

	TABLE '
Cationic Polymer	Anionio

	Cattorne Porymer	Anionic Micro-	rines first Pass
40	lbs/Ton	_bead lbs/Ton	Retention in %
	2.0 - 10 AEIMAC/90 AMD	-0-	50.3
	2.0 - 10 AEIMAC/90 AMD	2.0 - Silica- 5 rm	55.3
	2.0 - 10 AETMAC/90 AMD	5.0 - Bentonite	55.8
45	2.0 - 10 AETMAC/90 AMD	0.5 - 30 AA/70 AMD/	59.2

EXAMPLE 6

The procedure of Example 1 is again followed except that alum is added to the stock immediately before the cationic polymer. The test furnish is alkaline stock containing 5.0 lbs/ton of cationic starch and 55 25% CaCQ3. The results are set forth below in Table VI.

TA	RI	7	VI

Cationic Polymer lbs/Ton	Anionic Micro- bead-lbs/ton	Drainage in Seconds
	ton Alum	
0.5 - 10 AETMAC/90 AMD	5 - Bentonite	46.1
0.5 - 10 AETMAC/90 AMD	0.5 - 30 AMD/	39.9
	349 ppm MBA -130 rm	
10 lbs/t	ton Alum	

1 - 10 AEIMAC/90 AMD	5 - Bentonite -	33.5
1 - 10 AEIMAC/90 AMD	0.5 - 30 AA/70 AMD/ 349 ppm - 130 rm	29.6

The alum-treated furnish which is contracted with the polymer microbead has a faster drainage rate than that treated with 10 times as much bentonite. In a comparative test using 0.5 lb of 10 AETMAC/90 AMD and 5.0 lbs bentonite without alum, an equivalent drainage time of 46.1 seconds, is achieved.

25 EXAMPLE 7

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This example demonstrates the greater efficiency of the anionic organic polymer microbeads of the present invention used with alum as compared to bentonite alone. This efficiency is not only attained using a significantly lower anionic microbead dose but, also enable the use of a lower amount of cationic polymer. The furnish is alkaline and contains 5.0 libs/ton of cationic starch. The procedure of Example 1 is again used. The results are shown in Table VIII, bellow.

TABLE VII

	Cationic Polymer	Alum* 1bs/ton -0-	Anionic Microbead <u>lbs/ton</u> -0-	Drainage in Seconds 103.4
0	0.5 - 10 AETMAC/90 AMD	-0-	-0-	87.5
	0.5 - 10 AETMAC/90 AMD	5	-0-	
	0.5 - 10 AETMAC/90 AMD		-	76.4
	0.5 - 10 AETMAC/90 AMD	5	0.25 - 30 AA/ 70 AMD/349 ppm MBA	51.1
5			AND MEN SEE MEN	
	0.5 - 10 AETMAC/90 AMD	5	-130 mm 0.50 - 30 AA/70 AM	D 40.6
			349 ppm MBA-13 rm	
2	0.5 - 10 AETMAC/90 AMD	-0-	5 - Bentonite	51.6
	1.0 - 10 AETMAC/90 AMD	-0-	5 - Bentonite	40.2

* Alum is added immediately before the cationic polymer.

Thus, at a 0.5 lb. cationic polymer addition level, the anionic organic microbeads used with alum are

approximately 20 fold more efficient than bentonite used alone (0.25 lb. vs. 5.0 lbs.). The cationic polymer level can be reduced in half (0.50 lb. vs. 1.0 lb.) compared to bentonite when the microbead level is raised to 0.50 lb., which is 10 fold lower than the bentonite dose.

5 EXAMPLE 8

The procedure of Example 7 is again followed except that polyaluminum chloride is used in place of alum. As can be seen, in Table VIII, equivalent results are achieved.

10	TABLE VIII			
15	Cationic Polymer lbs/Ton 0.5 - 10 AEIMAC/90 AMD 0.5 - 10 AEIMAC/90 AMD	Aluminum Salt <u>lbs/Ton</u> -0- 5-Alum	Anionic Micro bead lbs/Ton Bentonite 0.5 - 30 AA/ 70 AMD/349 ppm	Drainage In Seconds 57.5 41.5
20	0.5 - 10 AEIMAC/90 AMD	8.5 Poly- aluminum Chloride (5.0 lbs alum (equivalent)	-130 rm 0.5 - 30 AA/ 70 AMD/349 pym -130 rm	42.0

EXAMPLE 9

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To a batch of alkaline paper stock is added cationic starch. The drainage time is measured after 30 addition of the following additives set forth in Table IX, below. The procedure of Example 1 is again used.

TABLE IX

35	Cationic Polymer 1bs/Ton	Anionic Microbead lbs/Ton	(Sec.) 5.0 lbs/Ton Starch	(Sec.) 10 lbs/Ton
40	0.5 - 10 AEIMAC/90 AMD 0.5 - 10 AEIMAC/90 AMD plus 5 lbs Alum	5 - Bentonite 0.5 - 30 AA/ 70 AMD/349 ppm MBA - 130 rm	46.9 34.0	50.9 32.7
	C - Comparation Short			

45 The alum/polymer microbead combination gives better drainage rates than the polymer/bentonite combination without alum.

EXAMPLE 10

First pass retention is measured on an alkaline furnish containing 5.0 lbs/fton of starch to which the additives of Table X, below, are added.

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TABLE X

Microbead 1bs/Ton 5.0 - Bentonite *5.0 - 30 AA/70 AMD/349 ppm MEA -130 rm	Fines First Pass Retention 10 AFIMAC/90 AMD (1bs/Ton) 0.5 1.0 2.0 39.9\$ 41.6\$ 46.8\$ 39.9\$ 44.4\$ 48.5\$
*5.0 - 30 AA/70 AMD/340	39.98 41.68 46.88 39.99 41.68

The microbead and bentonite give similar retentions with 0.5 lb/ton of cationic polymer but with higher concentrations of polymer better retention is obtained with the microbeads.

*With the anionic polymer microbead 5.0 lbs./ton of alum is added with the cationic polymer.

EXAMPLE 11

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Another alkaline paper furnish containing 5 lbs/ton of cationic starch and 2.5 lbs/ton of alum to which the additives of Table XI are added as in Example 10, is treated.

TABLE XI

30	TABLE_XT			
35	Anionic Microbead 1bs/Ton 5 - Bentonite 7 - Bentonite 0.25 - 30 AA/70 AMD/ 349 Dym MBA - 130 rm 0.5 - 30 AA/70 AMD/ 349 Dym MBA - 130 rm	Fines First 1 10 AEMAC/90 0.5 34.6 35.7 38.7	Pass Retention AMD (11bs/Ton) 1.0 42.38 43.18 43.48	

A significant reduction in the dosages of polymeric microbead results in equivalent or superior retention

EXAMPLE 12

Lower molecular weight, cationic, non-acrylamide based polymers are used in papermaking and in this example the effect of anionic microbeads on the performance of a polyamine of said class is set forth. To an alkaline furnish containing 5 lbs/ton of cationic, starch is added 1.0 lb/ton of a cationic polymeric polymer of 50 mole % epichlorohydnin, 47 mole % dimethylamine and 3.0 mole % ethylenediamine of 250,000 mol. 50 wt. The polyamine is used alone and in combination with 0.5 lbs/fon of microbead copolymer of 60% acrylic acid and 40% acrylamide cross linked with 1,381 ppm of methylenebisacrylamide and having 120 nm diameter particle size. From the data of Table XII it is seen that addition of the highly effective organic microbead cuts drainage time in half from 128.1 to 64.2 seconds.

TABLE XII

Cationic Polymer 1bs/Ton -0-	Anionic Microbead 1bs/Ton -0-	Drainage In Seconds 138.8
-1-	-0-	128.1
-1-	0.5	64.2

EXAMPLE 13

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In order to evaluate the use of microbeads on mill stock, a test is run on stock from a commercial paper mill. The paper stock consists of 40% hardwood/30 % soft wood/30% proke containing 12% calcium carbonate, 4% clay, and 2.5 lbs/hon of allyl succinic anhydride (ASA) synthetic size emulsified with 10 libston cationic potate starch. An additional 6 lbs/hon of cationic potate starch and 6 lbs/hon of tallum are also added to this stock. The additives listed in Table XIII, below, are added and drainage times are measured, as in Examble 1.

TABLE XIII

25	Cationic Polymer	Anionic Microbead 1bs/Ton -0-	Drainage In Seconds 153.7
	0.5 - 10 AEIMAC/90 AMD	-0-	112.8
30			
	0.5 - 10 AFTMAC/90 AMD	5.0 - Bentonite	80.3
	0.5 - 10 AETMAC/90 AMD	0.25 - 30 AA/	69.6
35		70 AMD -349 ppm	
		MBA - 130 rm	
	0.5 - 10 AEIMAC/90 AMD	0.5 - 30 AA/	57.5
		70 AMD - 349 ppm	
40		MBA - 130 mm	
	1.0 - 10 AEIMAC/90 AMD	5.0 - Bentonite	71.9
	1.0 - 10 AETMAC/90 AMD	0.5 - 30 AA/	49.1
45		70 AMD - 349 ppm	
		MPA - 130 mm	

The paper stock from the above run has a 153.7 second drainage time. Significant reduction of drainage time to 80.3 seconds is achieved with 0.5 lb/ton of high molecular weight, cationic polymer and 5 lbs/ton of bentonite. Replacement of the bentonite with a mere 0.25 lb/ton of organic anionic microbeads reduces drainage time another 10.7 seconds to 69.9 seconds. Thus, the microbeads at 1/20 the concentration give a superior drainage time to bentonite. The use of 0.5 lib/ton of organic above the drainage time to 57.5 seconds. This is 22.8 seconds taster than ten times the weight of bentonite.

When testing is carried out using 1.0 lb/ton of cationic polymer and 5.0 lbs/ton of bentonite, drainage time is 71.9 seconds. However, when the test is performed with 0.5 lb of microbeads, the drainage time is 49.1 seconds which is 22.8 seconds faster than bentonite with one tenth the amount of microbead.

EXAMPLE 14

The effect of using a cationic polymer of a lower charge density is investigated on the paper stock that was used in proceeding Example 13 and shown in Table XIV. The cationic polymer used, 5 AETMAC/95 5 AMD, has one half the charge density as that of 10 AETMAC/90 AMD that was used in Example 13. All else remains the same.

TABLE XIV

10	10			-
10	Cobinst - n s	Additional		
	Cationic Polymer	Alum*	Microbead	Drainage In
	0.5 - 5 AETMAC/95 AMD	lbs/Ton		_Seconds
		-0-	-0-	94.7
15	0.5 - 5 AEIMAC/95 AMD	-0-	5 - Bentonite	51.4
	0.5 - 5 AFIMAC/95 AMD	2.5	5 - Bentonite	
	0.5 - 5 AFIMAC/95 AMD			56.7
		-0-	0.5 - 30 AA/70 AMD/	48.7
20			349 ppm MBA-130 rm	
	0.5 - 5 AETMAC/95 AMD			
	AETHAC/95 AMD	2.5	0.5 - 30 AA/	39.5
			70 AMD/349	
25			ppm MBA -130 mm	•
25			Now and a Trick Limit	

- * Alum is added immediately before the cationic polymer.
- 30 The superiority of 1/10th the amount of polymeric microbead to bentonite is evident with a lower charge cationic polymer also. Furthermore, the drainage time of cationic polymer and bentonite did not improve but decreased by 5.3 sec. on further addition of 2.5 lbs/nor of alum.

EXAMPLE 15

The effect of changing the amount of starch on drainage time is measured by not incorporating the 6.0 lbs/non of additional starch added to the furnish in Example 13 using the same stock . The results are shown in Table XV.

••	*	TABLE XV		
45	Cationic Polymer	Additional Alum* lbs/Ton -0-		ninage In Seconds 45.9
50	0.5 - 5 AETMAC/95 AMD	-0-	0.5 - 30 AA/70 AMD/ 349 ppm MBA - 130 mm	39.5
	0.5 - 5 AETMAC/95 AMD	-2.5	0.5 - 30 AA/70 AMD/ 349 ppm MEA - 130 mm	29.5

* Alum is added immediately before the cationic polymer.

To evaluate the effect of the charge density of the cationic polymer on retention, to the furnish of Example 13, are added the additives shown in Table XVI. First pass retention values are measured, as in Example 5.

TABLE XVI

10	Alum* lbs/Ton -0-	Microbead 10s/Ton -0-	10 AFIMAC/90 AMD 0.5 lbs/Ton % Retention 36%	5 AETMAC/95 AMD 0.5 lbs/Ton % Retention 30.9%
	-0-	5 - Bentonite	32.4%	39.6%
	2.5	0.5 - 30 AA/70 AMD/	45.1%	49.1%

Table XVII (Cont'd)

349 ppm MBA -130 rm

			at 1.0 lbs/Ton	at 1.0 lbs/Ton
			% Retention	% Retention
25	-0-	5 - Bentonite	45.1	42.5
20	2.5	0.5 - 30 AA/70 AMD/ 349 ppm MBA - 130 rm	51.3 1	57.1

* Alum is added immediately before the cationic polymer.

Polymer microbeads are shown to be effective when used with high molecular weight, cationic polymers of lower charge density.

EXAMPLE 17

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A stock is taken from a second commercial mill. It is a goal of this example to demonstrate that 40 microbeads/atum give equivalent drainage times to those of current commercial systems. The mill stock consists of 45% delinked secondary fiber/25% softwood/30% broke containing 15% calcium carbonate and 3.0 lbs/ton of alkyl ketene dimer synthetic size emulsified with 10 lbs/fon of cationic starch. A second portion of 10 lbs of cationic starch is added to the thick stock and the ingredients listed in Table XVIII, below are added to the furnish, as described in Example 1.

TABLE XVII

Cationic Polymer lbs/Ton 0.6 10 AETAC/90AMD	Alum* lbs/Ton -0-	Anionic Microbead 1bs/Ton 5 - Bentonite	Drainage In Seconds 158.2 sec.
0.6 10 AFIMAC/90 AMD	-5.0	0.5 - 30 AA/70 AMD/ 349 ppm MBA -130 nm	141.6 sec.

* Alum is added immediately before the cationic polymer.

The microbeads/alum gives a faster drainage rate than the commercial bentonite system used in the mills routine production of paper. Other experimental runs result in lesser conclusive effectiveness with this pulp.

20 EXAMPLE 18

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Microbead retention efficiency is evaluated on papers made using a pilot Fourdrinier papermaking machine. The paper stock consists of pulp made from 70% hardwood and 30% softwood containing 25% calcium carbonate and 5 lbs/ton of cationic starch. The additives in the Table XVIII, below, are placed into 25 the furnish in successive runs and first pass retention percentages are measured. A 46 lb base weight paper is made.

The cationic, high molecular weight polymer is added just before the fan pump, the anionic microbead is added just before the pressure screen and alum, when added, is added just before the cationic polymer. Results are set forth in Table XVIII, below.

TABLE XVIII

5	Cationic Polymer lbs/Ton -0-	Alum lbs/Ton	Anionic Microbead lbs/Ton	Ash-Pirst Retention
	0.6 - 10 AEIMAC/90 AMD	-0-	7.0 - Bentonite	34.4% 61.3%
	0.6 - 10 AETMAC/90 AMD	2.5	0.25 - 30 AA/	62.7%
9			70 AMD/349 ppm MBA - 150 rm SV-1.32	
5	0.6 - 10 AETMAC/90 AMD	2.5	0.50 - 30 AA/ 70 AMD/349 ppm MBA - 150 mm	67.0%
			SV-1.32	

In this example, the combination of 0.5 lb/ton of microbeads and 2.5 lbs/ton of alum results in a 5.7% souperfor retention over 7.0 lbs/ton of bentonite alone. The 7.0 lbs/ton of bentonite is about equal to the combination of 0.25 lbs of beads and 2.5 lbs/ton of alum in retention properties, a significant dosage reduction.

EXAMPLE 19

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The same pilot paper machine and paper stock that was used in Example 18 is again used except that a 55 ib "basis weight" paper is made. Additives in Table XIX, below, are mixed into the furnish as in the preceding example on successive runs and retention values are measured.

TABLE XIX

5	Cationic Polymer lbs/Ton -0-	Alum 1bs/Ton -0-	Anionic Microbead 1bs/Ton -0-	Ash-First Pass Retention \$ 39.3%
	0.6 - 10 AEIMAC/90 AMD	-0-	-0-	39.4%
10	0.6 - 10 AEIMAC/90 AMD	-0-	7.0 Bentonite	74.6%
	0.6 - 10 AETMAC/90 AMD	2.5	0.5 - 30 AA/ 70 AMD/349 ppm	74.5%
15			MBA - 150 mm SV-1.32	
20	0.6 - 10 AFIMAC/90 AMD	5.0	0.5 - 30 AA/ 70 AMD/349 ppm	74.7%
26			MBA - 150 mm SV-1.32	

In comparing the heavier (55 lb) basis weight paper of Example 19 to that of Example 18 (46 lb), under all conditions, the heavier paper has better retention. With the heavier paper there is no significant difference in retention between the paper prepared with bentonite alone and that prepared with microbeads and either 2.5 lbs or 5 lbs of alum, except the significant dosage reduction i.e. 7lbs. vs. 0.5 lb.

EXAMPLE 20

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The effect of microbead on paper formation is evaluated by treatment of an alkaline furnish containing 35 5.0 lbs/ton of starch with the additives listed in Table XX, below, as described in Example 18.

TABLE XX

			Anionic	Paprican **
5	Cationic Polymer	Alum	Microbead	Microscanner
	lbs/Ton	1bs/Ton	lbs/Ton	SP/RMS Ratio
	1 - 10 AEIMAC/90 AMD	-0-	5 - Bentonite	66
0	1 - 10 AETMAC/90 AMD	-0-	1 - 30AA/70 AM 349ppm MBA- 130	

*Paper formation is measured on hand sheets in the Paprican micro-scanner as described by R. H. Trepanier, Tappi Journal, December pg. 153, 1989. The results indicate that the microbead treated paper has better formation at a lower dosage than the bentonite treated paper as the larger number signifies better formation.

EXAMPLE 21

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Using the paper stock of Example 20, except that the cationic starch concentration is increased to 10 lbs/ton, formation is measured on paper made with the additives set forth in Table XXI.

		Drainage Sec.	38.9	33.5
10		an nner <u>Ratio</u>		
15		Paprican Microscanner SP/RMS Ratio 73	81	77
20				
25	X		/40 AMD/	1/40 AMD/
30	TABLE XXI	Anionic Microbead 1bs/Ton	0.5 - 60 AA/40 AMD/ 1,381 ppm MBA	1.0 - 60 AA/40 AMD/ 1,381 ppm MBA
35) tr	. 04	
40		9	AMD .	AMD
45		Cationic Polymer 1bs/Ton	1 - 55 AETWAC/45 AMD	1 - 55 AETWAC/45 AMD
50		Cationic Polymer lbs/Ton	1 - 55	1 - 55

Microbeads give superior hand sheet paper formation and better drainage times compared to bentonite, and ss at a lower dosage.

EXAMPLE 22

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To an alkaline furnish containing 5-lbs of cationic starch, the ingredients set forth in Table XXII are added to the furnish of Example 21 and formation is observed visually on the paper hand sheets, produced

		TAUT SHARE		
tionic Lymer 12/Ton 1.5 - 10 AETWAC/90 AWD	Alum* lbs/ron	Anionic Microbead lbs/Ton	Visual Formation	Drainage
.5 - 10 AETMAC/90 AWD	5	-0-		87.8
Ser of Chiman of - 5.		5 - Bentonite	¥	57,5
OWN OF COURTS OF	2.5	0.5 - 30 AA/70 AMD/	ď	47.8
of the contract of the contrac		349 ppm MBA -130 nm		
to - 10 ASTRAC/90 AMD	ļ	5.0 - Bentonite	æ	49.2
TO ABIMAC/90 AMD	2.5	0.5 - 30 AA/70 AMD/	m	39.8
		349 ppm MBA - 130 nm		
1um 18 added immediately hefore the cationia	hefore the and			

* Alum is added immediately before the cationic polymer.

Hand sheets from the first three samples have equivalent formation (A) by visual observation. The last

two samples (B) themselves have equivalent formation by visual observation but their formation is not as good as the first three sheets. The experiment shows the superior drainage times are achieved with a microbead alum combination with equivalent visual paper formation as compared to bentonite, above, at higher dosage.

EXAMPLE 23

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In order to evaluate a different type of anionic microparticle, three different particle sizes of hydrophobic polystyrene microbeads, stabilized by sulfate charges, are added to an alkaline paper stock containing 25% to GCO₃ and 5 lbs/hon of cationic starch in the furnish. Table XXIII sets forth the additives used and drainage times measured.

TABLE XXIII

		Anionic	
15	Cationic	Polystyrene	
	Polymer	Microbeads	Drainage
	lbs/Ton	lbs/Ton	Sec.
	-0-		103.9 Sec.
	1.0 - 10 AETMAC/90 AMD	-0-	91.6 Sec.
20	1.0 - 10 AEIMAC/90 AMD	5.0 - Polystyrene beads- 98 mm	79.8 Sec.
	1.0 - 10 AEIMAC/90 AMD	5.0 - Polystyrene beads - 30 mm	49.9 Sec.
25	1.0 - 10 AETMAC/90 AMD	5.0 - Polystyrene beads - 22 mm	42.2 Sec.

It is noted that all three anionic polystryene microbeads improved drainage time over the cationic polymer alone with the smallest bead being the most effective.

The results indicate that noncross-linked, polymeric, water-insoluble microbeads are effective in increasing drainage rates.

EXAMPLE 24

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A 30 nm polystyrene bead is compared to bentonite in performance using the alkaline paper stock containing 5.0 libe/ton of cationic starch, above described in Example 22. Results are set forth in Table XXIV.

TABLE XXIV

	Cationic	Anionic	
	Polymer	Microbead	Drainage
	lbs/Ton	lbs/Ton	_Sec.
	1.0 - 10 AETMAC/90 AMD	-0-	70.9 Sec.
45	1.0 - 10 AFIMAC/90 AMD	5.0 - Bentonite	28.5 Sec.
	1.0 - 10 AEIMAC/90 AMD	5.0 - Polystyrene Beads - 30rm	30.5 Sec.

50 The results indicate that the 30nm polystyrene is substantially equivalent to bentonite.

EXAMPLE 25

Microbead size of anionic polymer is studied by measuring drainage rates on the alkaline paper stock of Example 23 to which the additives of Table XXV are added. Results are specified therein.

TABLE XXV

		TARREST STATES	
	Cationic Polymer	Anionic Microbead	
5	lbs/Ton		Drainage
	1.0 - 10 AETMAC/90 AMD	lbs/Ton	Sec
	1.0 - 10 AETMAC/90 AMD		106.8 Sec.
	AM	0.5 - 30 AA/70 AMD/	72.2 Sec.
	1.0 - 10 AFTMAC/90 AMD	349 ppm BMA - 130 rm	
10		2.0 - 40 AA/60 MBA	71.7 Sec.
	1.0 - 10 AETMAC/90 AMD	-220 nm	
	TO ALIMACY 90 AMD	0.5 - 30 AA/70 AMD/	98.9 Sec.
	1.0 - 10.37	50 ppm MBA - 1,000-2,000 r	מור
	1.0 - 10 AETMAC/90 AMD	2.0 - 30 AA/70 AMD/	102 E Com
15		50 ppm MBA- 1,000-2,000 rm	1

Both the 130 nm and 220 nm in diameter microbeads reduce drainage times over that of stock without microbeads by 33%. However, when the diameter of the anionic microbead is increased to 1,000 to 2,000 nm, drainage is not significantly effected.

EXAMPLE 26

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Using the same paper stock as in Example 22 the ingredients shown in Table XXVI are added in successive order, as in the previous examples. The results are specified.

TABLE XXVI

		TADDIC AAVI
30	Cationic Polymer <u>lbs/Ton</u> —0—	Anionic Microbeads Drainage 1bs/Ton Sec0- 135.6 Sec.
	1.0 - 55 AEIMAC/45 AMD	-0- 99.6 Sec.
35	1.0 - 55 AEIMAC/45 AMD	0.5 - 30 AA/70 AMD 86.7 Sec. 1000 ppm MEA- 2% surfactant
		-464 rm
40	1.0 - 55 AETMAC/45 AMD	0.5 lbs 30 AA/70 AMD/ 59.3 Sec. 1,000 ppm MEA- 4% surfactant
		-149 rm
45	1.0 - 55 AETMAC/45 AMD	0.5 lbs 30 AA/70 AMD/ 54.5 Sec. 1,000 ppm MEA-8% surfactant
		106 rm

Increased drainage rate is achieved as the microbead becomes smaller. Compared to the drainage time of 99.6 seconds without microbead without microbead at 99.6 seconds without microbead at 40% reduction, showing the effect of small diameter organic micropated or micropated at 40% reduction, showing the effect of small diameter organic micropated

EXAMPLE 27

To the same stock that was used in Example 23, the ingredients set forth in Table XXVII are added, as in said example.

TABLE XXVII

5	Cationic Polymer <u>lbs/Ton</u> 1.0 - 10 AETMAC/90 AMD	Anionic Microbeads <u>lbs/Ton</u> 0.5 - 30 AA/70 AMD/ 349 ppm MEA - 130 rm	Drainage <u>Sec.</u> 66.3
10	1.0 - 10 AETMAC/90 AMD	0.5 - 30 APS/70 AMD/ 995 ppm MBA SV-1.37 mPa.s	67.0

The microbeads of the 30 AA70 AMD/349 ppm MBA copolymer and those of the 30 APS/70 AMD/985 ppm MBA copolymer when used with cationic polymers, produces paper with aimset identical drained times, even though one has a carboxylate and the other has a sulfonate functional group. That the anionic beads have different chemical compositions and a differing degree of cross-inlining yet yield similar properties is attributed to this similar charge densities and similar particle size. The acrylic acid microbead to has a diameter of 130 nm and the 2-acrylamido-2-methyl-propane sulfonic acid microbead is of a similar size due to the similar way it was made.

EXAMPLE 28

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The effect of different shear conditions on the relative performance of the anionic microbead compared to bentonite is shown in Tables XXVIII & & B. Drainage testing is carried out as described in Example 1, on an alkaline furnish containing 5.0 lbs. of calcition: starch subjected to four different shear conditions.

TABLE XXVIII-A

		Stirring R.P.M. and Time*		
	Condition	Cationic Polymer Microbead		
	A	800 rpm-30 sec. 800 rpm-30 sec.		
35	В	1,500 rpm-30 sec. 800 rpm-30 sec.		
	С	1,500 rpm-60 sec. 800 rpm-30 sec.		
	D	1,500 rpm-60 sec. 1,500 rpm-5 sec		

High molecular weight cationic polymer is added to the furnish in a vaned Britt jar under spitation and spitation is continuous for the period specified before the microbead is added as in Example 1, agitation is continued, and the drainage measurement taken.

TABLE XXVIII-B

5	Cationic Polymer 0.6 lbs. 10 AETMAC/90 AMD	Anionic Microbead 5.0 lbs. Bentonite	Drainage in Seconds Shear Conditions A B C D 52.6 56.1 57.8 49.6
15	0.6 lbs.* 10 AETMAC/90 AMD	0.5 lbs. 30AA/ 70 AMD-349 ppm MBA-130 nm.	45.9 48.3 52.3 44.5

* 5.0 lbs. of alum is added immediately before the cationic polymer.

The relative performance of each additive system remains the same under different test shear conditions.

EXAMPLE 29

The utility of polymeric anionic microbeads in acid paper stock is established as follows. To an acid paper stock made from 2/3 chemical pulp 1/3 ground wood fiber, and containing 15% clay and 10 lbs/ton of alum at a pH of 4.5 are added and the listed ingredients of Table XXIV below.

35		TABLE XXIX	
40	Anionic Microbead 1bs/Ton -0- 5.0 - Bentonits 0.5 - 30 AA 70 AMD/ 349 Ppm MEA - 130 rm 1.0 - 30 AA/70 AMD/	Drainage using Cationic Polymer 10 AFIMAC/90 AMD 0.5 lbs/Ton 64.2 Sec. 57.0 Sec. 53.3	Drainage using Cationic Polymer 10 AETMAC/90 AMD 1.0 lbs/Ton 52.2 Sec. 47.0 Sec. 42.1 Sec.
	349 ppm MBA - 130 rm		38.7 Sec.

Thus, in acid paper processes,0.5 lb of polymeric anionic microbeads is superior to 5.0 lbs of bentionite in increasing drainage. At a level of 1.0 lbs/hon of cationic polymer, 5.0 lb/hon of bentionite lowers drainage time 10% while 0.5 lb/hon of microbeads lowers it 19.3% and 1.0 lb/hon of microbeads lowers it 25.9%.

EXAMPLE 30

This example demonstrates the effect of alum on drainage in the acid paper process when acid stock from Example 29 is used without initial alum addition. A set of drainage times is measured for this stock without alum present and a second series is measured with 5.0 lbs/cho or added atum and with lingredients set forth in Table XXX. The enhancement of drainage time with the added alum is a significant advantage of the present invention.

TABLE XXX

		Anionic	Drainage	in Seconds
	Cationic Polymer	Microbead	Alum in Stock	
5	lbs/Ton	lbs/Ton	-0-	5 lbs/Ton
	1.0 - 10 AETMAC/	5.0 - Bentonite	43.0	43.5
	90 AMD 1.0 - 55 AETMAC/	1.0 - 30 AA/70	42.1	29.1
10	45 AMD	AMD/ 349 ppm MBA - 130 rm	*	

C = Comparative Test

15 EXAMPLE 31

In recent years cationic potato starch and silica have been found to give improved drainage times when used in alkaline papermaking processes. The effectiveness of polymeric microbeads compared to the silica system is shown in Table XXXI using the ingredients set forth therein on to the alkaline paper stock of, and in accordance with, Example 1.

TABLE XXXI

25	Cationic Potato Starch lbs/Ton	Alum* lbs/Ton	Anionic Microbead lbs/Ton	Drainage Seconds
	-0-	-0-	-0-	119.1
	15 - Starch	-0-	-0-	112.7
	15 - Starch	5.0	-0-	84.3
30	15 - Starch	5.0	3.0 - Silica-5 mm	38.5
	15 - Starch	5.0	1.0 - 30 AA/70 AMD/	36.7
			349 ppm MBA-130 rm	
	30 - Starch	-0-	3.0 - Silica-5 rm	46.3

* Alum is added immediately before the addition of cationic potato starch.

The addition of 15 lbs/ton of starch, 5 lbs/ton of Alum and 3.0 lbs/ton of silica reduces the drainage time 67.7%, however replacement of the silica with 1.0 lb/ton of organic anionic microbeads reduces the drainage time 69.2% which is slightly better than the silica system with far less added material.

EXAMPLE 32

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The polymeric, anionic microbead and the silica starch systems of Example 31 are compared for first pass retention values using the alkaline paper stock of Example 2. The results are shown in Table XXXII, hellow.

TABLE XXXII

	Cationic Potato	THOUS WALL			
5	Starch 1bs/Ton -0-	Alum* 1bs/Ton -0-	Anionic Microparticle <u>lbs/Ton</u>	First Pass Retention	
	15 - Starch	-0-	3.0 - Silica 5 mm	25 %	
	15 - Starch	2 -		31.7 %	
10		2.5	0.5 - 30 AA/70 AMD/	37.4%	
	15 - Starch		349 ppm MBA- 130 mm		
	13 - Starch	2.5	1.0 - 30 AA/70 AMD/	46.6%	
			349 ppm MBA - 130 rm		
15					

- * Alum is added immediately before the addition of cationic potato starch.
- The retention values of starch and 3.0 lbs/non of silica are surpassed by replacing the silica with 2.5 lbs/non alum and either 0.5 lbs/non of microbead or 1.0 lb/non of microbeads. The process of the instant invention results in a 15.25% and a 34.1% improvement in retention values, respectively, over silica.

EXAMPLE 33

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Retention values using silica and the organic anionic microbead of Table XXXIII are compared in a pilot Fourdrinier papermaking machine. The paper stock consists of pulp made from 70% hardwood and 30% softwood containing 25% calcium carbonate and 5 isohon of cationic starch. The cationic potato starch is added immediately before the fan pump. The anionic microbeads and alum are added as in Example 18.

TABLE XXXIII

	Cationic Potato	Anom	VVVIII	
35	Starch 1bs/Ten -0- 20	Alum 1bs/Ton -0- -0-	Anionic Microbead 1bs/Ton -0- 3.0 - Silica 5 mm	Ash Retention
40	20 20	5.0 5.0	3.0 - Silica 5 rm 1.0 - 30 AA/70 AMD 349 ppm MBA - 150 rm SV-1.32	49.2 66.3% 68.7%

Alum improves the retention values of silica and the alum/silica system retention of 66.3% is slightly ses than that of the alum/organic anionic microbead system of 88.7% (3.5% improvement) with 1/3 the

EXAMPLE 34

A comparison of drainage times between the anionic, organic, microbead system and the silica system is made using the paper stock described in Example 13. It is noted that this stock contains 16 lbs/ton of cationic pot

TABLE XXXIV

Cationic Potato Starch	Alum** 1bs/Ton -00- 2.5	Anionic Microparticle 1bs/Ton 3.0 - Silica 5 rm 3.0 - Silica 5 rm 1.0 - 30 AA/70 AMD/ 349 ppm MEA -130 rm	Drainage Seconds 42.5 55.6 28.7
------------------------	-------------------------	---	---

** Alum is added immediately before the addition of cationic potato starch.

(*Reverse addition of silica before starch)

The silica/starch system is inferior in drainage time to that of the organic microbead system (1.0 lb and 2.5 lbs alum).

EXAMPLE 35

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With the same stock as in Example 34, organic, anionic, microbead and silica systems, using a anionic polymer added to the furnish, are compared as to drainage times as in said Example. Alum and cationic starch are added where indicated and the furnish is stirred at 800 r.p.m. for 30 seconds. The anionic acrylamide copolymers and, if added, silica or microbeads are added together to the furnish and stirred for 25 a further 30 seconds at 800 r.p.m. before the drainage rate is measured. See Table XXXV.

TABLE XXXV

30	Anionic Polymer Retention Aid 1bs/Ton -0-	Alum* <u>lbs/Ton</u> -00-	Anionic Microbead 1bs/Ton -0-	Drainage Seconds 92.4 62.1
35	0.3 - 30 AA/70 AMD 0.3 - 30 AA/70 AMD 0.3 - 30 AA/70 AMD 0.3 - 30 AA/70 AMD 0.3 - 30 AA/70 AMD	5.0 -0- -0- 5.0	-0- 0.5 - Silica-5 rm 1.0 - Silica-5 rm 0.5 - 30 AA/70 AMD/ 349 ppm MEA - 130 rm	59.4 50.4 47.5 42.2
40	0.3 - 30 AA/70 AMD and 10 - additional cationic starch	-0-	1.0 - Silica-5 rm	
45	0.3 - 30 AA/70 AMD and 10 additional cationic starch	5.0	0.5 - 30 AA/70 AMD/ 349 ppm MBA-130 rm	28.4

* Alum is added immediately before the addition of cationic potato starch, where both one used.

Silica improves drainage times when compared to the anionic acrylamide polymer alone; however, the anionic organic microbeads, in replacing the silica, give even better drainage times with alum. Additional cationic potato starch in the furnish allows the microbead system to produce even faster drainage times.

EXAMPLE 36

Comparative retention values are determined for an organic anionic microbead versus a silica system using an anionic polymer and the paper stock of Example 13. The additives, as specified in Table XXXVI, are added as in Example 35.

TABLE XXXVI

5	Anionic Polymer	Alum 1bs/Ton -0-	Anionic Microbeadlbs/Ton	First Pass Retention 8 34.3
	0.3 - 30 AA/70 AMD	5.0	-0-	37.3
	0.3 - 30 AA/70 AMD	-0-	1.0 - Silica-5 rm	34.0
10	0.3 - 30 AA/70 AMD	-0-	0.5 - 30 AA/70 AMD/	40.3
			349 ppm MBA-130 rm	
	0.3 - 30 AA/70 AMD	5.0	0.5 - 30 AA/70 AMD	52.6
15			349 ppm MBA-130 rm	

Retention values with 0.3 lb/ton of anionic polymer, with and without silica, are identical at 34% and addition of 5.0 lbs/ton of alum and no silica actually increases retention to 37.3%.

Anionic polymers, in combination with organic anionic microbeads however, give better retention values without (40,3%) and with alum (62,8%) when compared to the silica system (34%). This retention when combined with the faster drainage rates of the organic anionic microbeads shown in Table XXXV, makes them preferable to either the silica or bentonite systems usually used commercially.

EXAMPLE 37

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The effect of cationic organic, microbeads is now examined. To an alkaline furnish containing 25% calcium carbonate, 15 lbs. of cationic starch and 5 lbs. of alum and of a pH of 8.0, the ingredients of Table XXXVII are added. The anionic polymer is added first and the cationic, organic microbead is added second.

	TABLE XXXVII
	Cationic

		Cattorne	
	Anionic Polymer	Microbead	
	(Linear)	or Polymer	Drainage
35	lbs/Ton	_lbs/Ton	Seconds
	-0-	-0-	142.7
	0.5 - 30 AA/70 AMD	-0-	118.5
	0.5 - 30 AA/70 AMD	0.5 - 40 AETMAC/60 AMD/	93.3
40		100 ppm MBA- 100 rm	
	0.5 - 30 AA/70 AMD	0.5 - 40 AEIMAC/60 AMD/	113.9
	•	100 ppm MBA - 1,000 mm	
45	0.5 - 30 AA/70 AMD	0.5 - 40 AEDMAC/60 AMD/	98.7
~		1 P-1	
		linear Polymer	
		(not a microbead)	
		•	

The addition of 0.5 lb/ton of cross-linked cationic microbead - 100 nm results a drainage time reduction of 25.2%. Addition of 0.5 lb/ton of linear cationic polymer causes a drainage time reduction but is not as effective as the cationic microbeads of the present invention.

EXAMPLE 38

To an acid paper stock made from 2/3 chemical pulp, 1/3 ground wood fiber and 15% clay are added 20 lbston of alum. Half the stock is adjusted to pH 4.5 and remainder is adjusted to pH 5.5. The Ingredients shown in Table XXXVIII are added in the same order as Example 37.

	ords ords H 5.5 15.5 55.8 59.7 74.7 74.7 92.8 55.0 62.8 55.4 62.8 55.4
5	Drainage Tithe Transcords 103.4 bt 15.5 bt 16.4 bt 16.
10	7,60 AND/ 3,760 AND/ 300 rm 2,60 AND/ 0 rm 000 rm 0,70 AND/ 10 mm 000 rm
15	Cationic Hicrobead Historic Alica Cationic Hicrobead Historic Control
20	00.5 00.5 100 100 100 100 100 100 100 100
25	Cottonic Polymer 1152/751 -000- 0.5 - 40 AETMAC/60 AND 0.5 - 40 AETMAC/60 AND -0000000000
30	Cattonio Polymer 125.779 -000000000
35	ċ ċ · · · ·
40	JAMES - 10 AES/90 AUD - 10 AES
45	Antionida 1960/2002 10.5 - 0

Examples 39-45

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Following the procedure of Example 2, various microbeads, high molecular weight (HMN) polymers and so polysaccharides are added to paper-making stock as described therein. In each instance, similar results are observed.

				HMW
5	Example No.	Microbead	Polysaccharide	Polymer
	39	AM/MAA (50/50)	Cationic Guar	AM/DADM (70/30)
10	40	AM/VSA (65/35)		Mannich PAM
15	41	Mannich PAM	CMC	AM/AA (80/20)
20	42	AM/DADM (75/25)		PAA
	43	P(DMAEA)		
25	44	P(AA)	Cationic Guar	AM/ DMAEA
30	45	AM/AA (25/75)	Cationic Guar	AM/AA (70/30)
35		rylamide ethacrylic acid		

MAA = Acrylamide

VSA = Vinyl Sulfonic acid

DADM = Diallydimethylammonium chloride

P(AA) = Polyacrylic acid
P(DMAEA) = Poly(dimethyler

P(DMAEA) = Poly(dimethylaminoethylacrylate) quaternary
CMC = Carboxymethyl cellulose

Mannich PAM = Polyacrylamide reacted with formaldehyde and diemthyl amine

Claims

- 45 1. A method of making paper which comprises adding to an aqueous paper furnish from about 0.05 to about 20 lbs/fon, based on the dry weight of the paper furnish solids, of an ionic, organic, polymeric microbead, the microbead being less than about 750 nanometers in diameter if cross-linked and less than 60 nanometers in diameter if non-cross-linked and water-insoluble, the ionicity of said microbead being at least 1%, but at least 5%, if cross-linked, anionic and used alone.
 - A method according to Claim 1 wherein from about 0.05 to about 20 lbs/ton, same basis, of a high molecular weight, ionic polymer is added to said furnish in conjunction with said microbead.
 - A method according to Claim 1 wherein from about 1.0 to about 50 lbs/ton, same basis, of an ionic polysaccharide is added to said furnish in conjunction with said microbead.
 - A method according to Claim 1 wherein from about 0.1 to about 20 pounds of an active, soluble aluminum species is also added per ton of paper furnish solids to the furnish.

- 5. A composition of matter comprising a mixture of A) an ionic, organic, polymer microbad being less than about 750 nanometers in diameter if cross-linked and less than about 60 nanometers in diameter if non-cross-linked and water-insoluble, the lonicity of the microbad being at lesst about 1% and either B) a high molecular weight lonic polymer, the ratio of AE ranging from about 1:400 to about 400:1, respectively, or C) in lonic polysaccharide, the ratio of AC ranging from about 20:1 to about 1:1000 or B and C together, the ratio if AEB and C together, the ratio if AEB and C together tranging from about 400:1 to about 1:1000.
- 6. A composition according to Claim 5 containing, in addition, an active, soluble aluminum species.
- 7. Paper produced by the method of Claim 1.
 - 8. A method according to Claim 1 wherein bentonite or silica is added in conjunction with the microbead.
 - 9. A composition according to Claim 5 containing, in addition, bentonite or sillca.
 - Paper produced by the method of Claim 8.

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EUROPEAN SEARCH REPORT

Application Number

EP 91 10 4837

Category	Citation of docume	nt with indication, where eppropri f relevant passages	ate,	Relevant to claim	CLASSIFICATION OF THE
х	EP-A-0 315 718 (SUM LTD.) * the whole document *	ITOMO CHEMICAL COM	PANY,	1,7	D 21 H 21/54
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